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(71) Applicant: Astra Aktiebolag
151 85 Södertälje (SE)

(72) Inventors:

- Bowald, Staffan
740 10 Almunge (SE)
- Johansson-Ruden, Gunilla
436 42 Askim (SE)

Remarks:

This application was filed on 24 - 07 - 1996 as a
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(54) **A novel surgical material**

(57) Polyhydroxybutyric acid polymers or copolymers of hydroxybutyric acid and hydroxyvaleric acid may be used for preparing a porous, bioresorbable flexible sheet for tissue separation and stimulation of tissue regeneration at healing processes in injured soft tissue in mammals including man. The sheet has a pore size permitting passage of water and salts through the sheet but which locks out cells and other tissue particles. Processes for preparation of the material and are also described.

Description**Technical Field**

The present invention is related to a material for separation of injured tissue at healing processes. It is further related to a process for preparation of such material, and to use of the material in healing processes.

Background of the Invention

In healing of injuries in soft tissue, scar tissue develops, which in many cases disturbs the function of the damaged organ and adjacent organs. This problem primarily occurs in healing of internal organs, while the problems in healing of skin and external mucosae may also be of cosmetic nature. Reference to injury and injured soft tissue herein primarily relates to incisions and other injuries caused by surgical operations in the organ subject to surgical correction, as well as in covering and adjacent organs. Among injuries caused by surgical operations is included injuries caused by surgical correction of congenital defects e.g. fistulas. The material according to the invention may also advantageously be employed in healing of injuries caused by external violence e.g. in accidents. A purpose of the invention is to facilitate and improve the healing by locking out undesired cells, other tissue and/or foreign particles. For this purpose, porous cloth of polytetrafluoroethylene such as Gore-Tex® is currently in use. The disadvantage with this is that foreign material will remain in the body, which may cause problems. Bowald *et al.* in *The Lancet* No. 8056, January 21, 1978, page 153 describes the use of a knitted mesh of polyglactin 910 (Vicryl®) as an arterial substitute. A preclotted mesh was sutured as a patch graft or as an end to en tube in the thoracic aorta in pigs. This coarse-mesh material is only useful by deposition of fibrin intermingled with platelets and red blood cells in the mesh spaces. Further studies on coarse-mesh polyglactin material were reported in *Surgery* vol. 86, No. 5, pp. 722-729, 1979, in *Scand. J. Thor. Cardiovasc. Surg.* 15:91-94, 1981, in *Muscle & Nerve* 5:54-57, 1982 and in *Acta Chir. Scand.* 146:391-395, 1980. SE 8604571-3 describes the use of resorbable and non resorbable membranes for accelerating bone formation and bone healing. However, cellular processes of resorptive type are indicated as undesirable, as they may delay bone formation and damage the newly formed bone.

US-A-4603695 (Ikada *et al.*/Japan Medical Supply Co., Ltd.) discloses a number of materials which may be used to prevent adhesion of vital tissues which are absorbable by degradation in vital tissues. Polymers and copolymers of lactic acid are preferred, though a polyester of β -hydroxybutylcarboxylic acid is also mentioned.

Biomaterials, vol. 8, 129-137, 1987 (Miller *et al.*) discloses a study on the biodegradation of poly- β -

hydroxybutyrate (PHB) homopolymer and poly- β -hydroxybutyrate-hydroxyvalerate copolymers. The conclusion drawn by the authors is that PHB is generally not biodegradable *in vivo*. Furthermore, they state that the biodegradation of implanted PHB and its copolymers cannot be predicted with confidence.

Description of the Invention

According to the invention it has now been made possible to avoid such problems as mentioned above in healing of soft tissue. The invention provides the use of polyhydroxybutyric acid polymers or copolymers of hydroxybutyric acid and hydroxyvaleric acid for preparing a porous, bioresorbable flexible sheet for tissue separation and stimulation of tissue regeneration at healing processes in injured soft tissue in mammals including man, said sheet having a pore size permitting passage of water and salts through the sheet but which locks out cells and other tissue particles. Using this material has been found to cause a specific stimulating effect on formation of macrophages in soft tissue. The macrophages release a growth factor which stimulates tissue healing. Using this material of the invention does not require preclotting or the presence of blood for functioning.

According to a preferred embodiment of the invention the material has a pore size up to 30 μm , preferentially 0.1-10 μm . The sheet thickness may be between 1 μm and 5 mm, but is preferably 10 μm to about 1 mm.

The sheet may be adapted to replace a part of the pericardium, but equally well it may not be adapted to replace a part of the pericardium.

The invention also provides a porous, bioresorbable flexible sheet for tissue separation and stimulation of tissue regeneration at healing processes in injured soft tissue in mammals including man, said sheet being made of polyhydroxybutyric acid polymers or copolymers of hydroxybutyric acid and hydroxyvaleric acid and having a pore size permitting passage of water and salts through the sheet but which locks out cells and other tissue particles.

The material may according to the invention be prepared according to the following processes, of which non-woven technique, precipitation and laser technique are preferred:

Non-woven

Non-woven fibrous material is prepared as described in US Patent No. 4 603 070. Fibres are produced from a melt or solution of the polymer by pressing the material through a perforated outlet. The fibres are spread randomly, or with a main orientation, on a support (a still glass plate—a mobile net ribbon—other mould). In this manner a porous "cloth" is obtained which may be given varying porosity by modification of fibre dimension, spreading method, material thickness and/or working up with heat/compression. The thick-

ness of the cloth is preferably 300—500 µm.

Perforation

A homogeneous film/cloth of the material may be perforated by e.g. laser technique to achieve porosity. In particular, a weak so-called excimer laser may be employed together with a template of perforated stainless steel.

Precipitation (not applicable to PGA)

The polymer is dissolved in a solvent which may be selected from a first group comprising dimethylformamide (DMF), dimethylacetamide (DMA), dimethyl sulfoxide (DMSO), and tetrahydrofuran (THF), or from a second group comprising chlorinated hydrocarbons such as chloroform and methylene chloride. Precipitation of the polymer may be achieved with a precipitation agent, which with the first group of solvents suitably is water, possibly with an addition of up to 20% solvent. With the second group of solvents ethanol and other lower alcohols may be used as a precipitation agent, possibly with an addition of up to 20% solvent. Both the solvent and the precipitation agent may be a mixture. Temperature and time at the precipitation may be selected to achieve any desired pore size.

Effervescence

By admixture of an effervescent agent which releases gas e.g. in contact with water (at precipitation) or on heating (in a melt).

Leaching

Soluble particles, for example salt, are suspended in a solution of the polymer/admixed into a melt thereof. After evaporation/solidification the particles are washed out of the material by leaching in a suitable solvent for the particles (but not for the polymer). This washing can be done completely, or partially provided that a non-toxic salt such as NaCl is employed, whereby the residual amount of salt may be allowed to leach out after implanting the material into the body.

The material is used according to the invention in healing of soft tissue, i.e. tissue that does not consist of cartilage, bone or teeth. Preferably the material is used for healing of injuries in the:

- Circulatory system (heart, blood vessels such as the pulmonary artery)
- Digestive organs (stomach, intestines, oral cavity, liver, pancreas)
- Reproductive organs (uterus e.g. in a Caesarean section, ovaries, testes e.g. at undescended testis in boys, the Fallopian tubes, testicular ducts)

- Urinary system (kidney, bladder, urethra)
- Respiratory system (lungs, trachea, bronchi)
- Other muscles (abdominal wall etc.)

Suitable bioresorbable materials for the purposes of the present invention may readily be chosen by one skilled in the art, e.g. among those that are either commercially available or have been described in literature or will be available in the future. As examples of such bioresorbable materials may be mentioned polymers based on polyglycolic acid (PGA), copolymers of glycolic acid and lactic acid, copolymers of lactic acid and ϵ -aminocapronic acid, and various lactide polymers. PGA esters are, e.g. described in US Patent No. 3 463 658, while copolymers of glycolic acid and lactic acid are described e.g. in US Patent No. 3 982 543. Homo and copolymers of lactic acid are described in e.g. US Patent No. 3 636 956. Examples of commercially available materials are Vicryl® (a copolymer of 90% glycolic acid sold by Ethicon, Sommerville, N.Y., USA—also known as Polyglactin) and Dexon® (Davies & Geck, Pearl River, N.Y., USA). Further examples are polydeshoxazone (PDS) (Ethicon, USA), polyhydroxybutyric acid (PHB), copolymers of hydroxybutyric acid and hydroxyvaleric acid (PHBV), polyesters of succinic acid, and crosslinked hyaluronic acid. As suggested above, mixtures of the above-mentioned materials may equally well be employed. One skilled in the art would have no difficulty to modify such bioresorbable materials depending on current needs, e.g. with regard to resorption time, strength etc.

Possibly, growth factors may be included in the porous structure, either deposited in the pores or included in the bioresorbable material for slow release of growth factor.

The sheet-formed material according to the invention may suitably, in particular in application for strong muscles and in other locations where the material is subject to strong load, be combined with a resorbable armament e.g. a woven or knitted cloth.

The invention is further described with reference to the following examples.

Example 1

Preparation of a sheet material for replacement of a part of the pericardium.

50 5 ml of a solution of 10 g Biopol (PHBV, 20% hydroxyvaleric acid) in 100 ml dimethylacetamide (about 50 °C) was spread on a glass plate. The glass plate was thereafter placed in water of ambient temperature for 12 hours. In this manner a porous patch (8 x 8 cm) was formed having about 1 mm thickness. The patch was washed in water, dried, packed and sterilized (ethylene oxide).

Example 2Use for healing of pericardiac defects.

In connection with cardiac surgery, difficulties occur almost always in closing the pericardium. This results in the pericardium often being left open. The result is adhesion which causes severe difficulties on re-operations and also an decreased motility of the heart, the function of which is impaired.

In connection with cardiac operations on sheep the defect caused was replaced with a patch of tissue-compatible resorbable polymer prepared according to Example 1. The patch was stitched into the defect by a continuous suture. When the animal after four months of healing was sacrificed and autopsy was performed, virtually normal pericardiac tissue was found to be formed without growing together with the heart surface, and the heart had been freely motile in the pericardium.

Example 3Producing a nonwoven patch for reconstruction of pericardium.

The nonwoven material was made from solution spun PHB-fibres pressed together to a patch (produced in accordance with US patent 4 603 070). Patch thickness was about 0.4 mm with about 70 per cent pore volume, patch size 15 x 15 cm. The patch was sterilized in ethylene oxide.

Example 4

Nonwoven PHB patches, produced according to Example 3, was used to replace a part of the pericardium in 10 sheep. The animals have been followed up for more than one year after the operation and have been sacrificed at different times. After two months regeneration of the pericardium had started, a very loose adhesion could be found. In the tissue a very active phagocytosis, with macrophages as the dominating type of cells, could be seen. No other kind of inflammation was present.

Later there were no signs of adhesion and already after four months a healing, very much like normal pericardium could be seen. The inner side was very smooth and glossy and mesothelial cells were present, which means that real pericardium had regenerated.

Up to ten months a slight darkness of the patch area could be observed due to partly remaining polymer. The darkness disappeared when all polymer was resorbed.

Example 5Producing a tube for urethra reconstruction.

Vicryl®-fibre was knitted to form a thin tube. The

tube mesh, 10 cm long, was mounted on a glass stick, diameter 4 mm. The tube was dipped in a solution of 10 g PHB:HV (80:20) in 100 ml DMAc and then dipped in water for 12 hours to get a porous structure. After washing, drying and packaging the urethra tube was sterilized in ethylene oxide.

Example 6

10 The urethra in 4 dogs was replaced by a urethra tube, produced according to Example 5. Six to nine months later the prosthesis had been resorbed and a fully functional urethra tissue was reconstructed in all animals.

Claims

- 15 1. Use of polyhydroxybutyric acid polymers or copolymers of hydroxybutyric acid and hydroxyvaleric acid for preparing a porous, bioresorbable flexible sheet for tissue separation and stimulation of tissue regeneration at healing processes in injured soft tissue in mammals including man, said sheet having a pore size permitting passage of water and salts through the sheet but which locks out cells and other tissue particles.
- 20 2. Use in accordance with claim 1, in which the pore size is up to 30 µm, preferentially 0.1—10 µm.
- 25 3. Use in accordance with claim 1 or 2, in which the thickness of the sheet is about 10 µm—1 mm.
- 30 4. Use accordance with any one of the preceding claims, said sheet being combined with a resorbable reinforcement.
- 35 5. Use accordance with any one of the preceding claims, provided that said sheet is not adapted to replace a part of the pericardium.
- 40 6. A porous, bioresorbable flexible sheet for tissue separation and stimulation of tissue regeneration at healing processes in injured soft tissue in mammals including man, said sheet being made of polyhydroxybutyric acid polymers or copolymers of hydroxybutyric acid and hydroxyvaleric acid and having a pore size permitting passage of water and salts through the sheet but which locks out cells and other tissue.
- 45 7. A sheet in accordance with claim 6, in which the pore size is up to 30 µm, preferentially 0.1—10 µm.
- 50 8. A sheet in accordance with claim 6 or 7, in which the thickness of the sheet is about 10 µm—1 mm.
- 55 9. A sheet in accordance with any one of claims 6 to 8, said sheet being combined with a resorbable rein-

forcement.

10. A sheet in accordance with any one of claims 6 to 9,
provided that said sheet is not adapted to replace a
part of the pericardium. 5

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EUROPEAN SEARCH REPORT

Application Number
EP 96 20 2092

DOCUMENTS CONSIDERED TO BE RELEVANT			CLASSIFICATION OF THE APPLICATION (Int.Cl.)		
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim			
D,X	US-A-4 603 695 (IKADA YOSHITO ET AL) 5 August 1986 * column 3, line 45 - line 63 * * column 4, line 33 - line 50 * ---	1-3,6-8, 10	A61L27/00 A61L31/00		
X	DATABASE WPI Section Ch, Week 8535 Derwent Publications Ltd., London, GB; Class A96, AN 85-214070 XP002018223 & JP-A-60 137 402 (MITSUBISHI RAYON KK) , 22 July 1985 * abstract * ---	1			
A	US-A-4 355 426 (MACGREGOR DAVID C) 26 October 1982 * claims *	1-10			
A	WO-A-86 02843 (WALLSTEN HANS IVAR) 22 May 1986 * claims; examples *	1-10			
D,A	US-A-4 603 070 (STEEL MARGARET L ET AL) 29 July 1986 * claims *	1-10	A61L		
A	DATABASE WPI Section Ch, Week 8608 Derwent Publications Ltd., London, GB; Class A88, AN 86-053312 XP002018224 & JP-A-61 008 107 (MITSUBISHI RAYON KK) , 14 January 1986 * abstract * ---	1-10			
A	EP-A-0 204 931 (MATERIALS CONSULTANTS OY) 17 December 1986 * claims; example 8 *	1-10			
		-/-			
The present search report has been drawn up for all claims					
Place of search	Date of completion of the search	Examiner			
THE HAGUE	11 November 1996	ESPINOSA, M			
CATEGORY OF CITED DOCUMENTS					
X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document					
T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document					



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EUROPEAN SEARCH REPORT

Application Number
EP 96 20 2092

DOCUMENTS CONSIDERED TO BE RELEVANT			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
A	FR-A-2 060 524 (WESOLOWSKI) 18 June 1971 * claims *	1	
A	WO-A-86 07250 (MATERIALS CONSULTANTS OY) 18 December 1986 * claims; examples *	1	
A	EP-A-0 258 781 (AMERICAN CYANAMID CO) 9 March 1988 * claims *	1	

			TECHNICAL FIELDS SEARCHED (Int.Cl.6)
	The present search report has been drawn up for all claims		
Place of search	Date of completion of the search	Examiner	
THE HAGUE	11 November 1996	ESPINOSA, M	
CATEGORY OF CITED DOCUMENTS		T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons ----- & : member of the same patent family, corresponding document	
X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document			